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=> s (tryptophan(w)hydroxylase(w)isoform)/bi.ab 69687
TRYPTOPHAN/B 54633 TRYPTOPHAN/AB
39674 HYDROXYLASE/B 34573 HYDROXYLASE/AB
43265 ISOFORM/B 28280 ISOFORM/AB
L1 15
(TRYPTOPHAN(W)HYDROXYLASE(W)ISOFORM)/BI,AB

=> s tph2/bi.ab 23 TPH2/B 23 TPH2/AB
L2 23 TPH2/BI,AB

=> s l1 or l2
L3 28 L1 OR L2

=> d his
(FILE 'HOME' ENTERED AT 15:54:17 ON 25 MAY 2005)
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L1 15 S
(TRYPTOPHAN(W)HYDROXYLASE(W)ISOFORM)/BI,AB
L2 23 S TPH2/BI,AB
L3 28 S L1 OR L2

=> d l3 1-28 bib ab

L3 ANSWER 1 OF 28 CAPLUS COPYRIGHT 2005 ACS on STN
AN 2005:405351 CAPLUS

TI Method for identifying compounds that affect expression of
 tryptophan ***hydroxylase*** ***isoform*** 2
 IN Clark, Janet
 PA Merck & Co., Inc., USA
 SO PCT Int. Appl., 59 pp. CODEN: PIXXD2

DT Patent
 LA English
 FA/CNT 1 PATENT NO. KIND DATE APPLICATION
 NO. DATE

PI WO 2005/041750 A2 20050512 WO 2004-US34619
 20041020 W. AE, AG, AL, AM, AT, AU, AZ, BA, BG, BR,
 BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ,
 EE, EG, ES, FI, GB, GR, GE, GH, GM, HR, HU, ID, IL,
 IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV,
 MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM,
 PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
 TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW,
 RW, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW,
 AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY,
 CZ, DE, DK, EE, ES, FI, FR, GB, GR, GU, IE, IT, LU, MC, NL,
 PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, GM, GA, GN,
 GQ, GW, ML, MR, NE, SN, TD, TG
 PRAI US 2003-514268 P 20031024

AB A method for identifying analytes which directly or indirectly
 affect ***tryptophan*** ***hydroxylase***

isoform 2 (***TH2***) expression is described.
 The method enables glucocorticoid receptor modulators and
 8βγ-hydroxysteroid dehydrogenase type 1 (8βγ-HSD1)
 inhibitors to be screened for central nervous system penetrance
 and activity by determining their ability to regulate expression of
 TH2. The method is particularly useful for identifying
 analytes which suppress glucocorticoid disruption of central
 serotonergic neurotransmission in the brain.

L3 ANSWER 2 OF 28 CAPLUS COUNTRY GHT 2005 ACS ON STN
 AN 2005.364511 CAPLUS

TI A second ***tryptophan*** ***hydroxylase***
 isoform TH-2 mRNA, is increased by ovarian steroids in
 the raphe region of macaques

AU Sanchez, Rachel L.; Reddy, Arubala P.; Centeno, Maria L.;
 Henderson, Jessica A.; Bethesda, Cynthia L.

CS Division of Reproductive Sciences, Oregon National Primate
 Research Center, Beaverton, OR, 97006, USA

SO Molecular Brain Research (2005), 135(1-2), 194-203

CODEN: MBREB4; ISSN: 0169-328X

PB Elsevier B.V.

DT Journal

LA English

AB Recently, a second gene that codes for the rate-limiting
 enzyme in serotonin synthesis was found in brain, named
 tryptophan hydroxylase-2 (TH-2). We sequenced overlapping
 segments (251 and 510 bp) of 5' monkey TH-2 and questioned
 whether TH-2 is regulated by estrogen (E) and progesterone (P)
 in serotonin neurons of macaques. Monkey TH-2 was 97%
 homologous to human TH-2 and 65% homologous to monkey
 TH-1 in the coding region. Spayed monkeys were administered
 placebo, E-only, P-only, or E + P for 1 mo via Silastic implants (n
 = 4/treatment) and the midbrain was utilized for TH-2 in situ
 hybridization (ISH). Admt. monkeys (n = 3/treatment) were
 used to def. the relative abundance of TH-2 mRNA with quant.
 (q) RT-PCR. In the ISH assay, all of the hormone treatments
 caused a significant and similar increase in TH-2 mRNA optical
 d. (fourfold; P < 0.004) and pos. pixel area (twofold; P < 0.002)
 over spayed controls. Treatment with E or E + P for 1 mo
 increased the relative abundance of TH-2 mRNA over spayed
 controls in the qRT-PCR assay (ANOVA P < 0.05 and P < 0.007,

resp.). In conclusion, ovarian steroids stimulate TH-2 mRNA
 expression, which could in turn cause an increase in serotonin
 synthesis. This would impact many of the neural functions that
 are governed by serotonin.

L3 ANSWER 3 OF 28 CAPLUS COUNTRY GHT 2005 ACS ON STN
 AN 2005.345675 CAPLUS

TI Promoter polymorphism of second ***tryptophan***
 hydroxylase ***isoform*** (***TH2***) in
 schizophrenia and suicidality

AU De Luca, Vincenzo; Volmekos, Daphne; Wong, Gregory W.
 H.; Shinkai, Takahiro; Rothe, Claudia; Strauss, John; Kennedy,
 James L.

CS Neurogenetics Section, Clarke Site, Centre for Addiction and
 Mental Health, Department of Psychiatry, 250 College Street,
 University of Toronto, P-30, Toronto, ON, M5T 1B8, Can.

SO Psychiatry Research (2005), 134(2), 195-198 CODEN:

PSRSDR; ISSN: 0165-1781

PB Elsevier Ltd.

DT Journal

LA English

AB Allele and haplotype frequencies of a promoter
 polymorphism in the gene encoding tryptophan hydroxylase (
 TH2) did not differ in 83 suicidal schizophrenic patients
 compared with 170 non-suicidal schizophrenic patients. These
 findings suggest that these 5' marker haplotypes in the
 TH2 gene do not influence suicidal behavior in
 schizophrenia.

L3 ANSWER 4 OF 28 CAPLUS COUNTRY GHT 2005 ACS ON STN
 AN 2005.309053 CAPLUS

TI Differential hormonal regulation of tryptophan hydroxylase-2
 mRNA in the murine dorsal raphe nucleus

AU Clark, Janet A.; Pai, Lee-Yuh; Rick, Rosemarie Beth; Rohrer,
 Susan P.

CS Department of Molecular Endocrinology, Merck Research
 Laboratories, Merck & Company Inc., Rahway, NJ, USA

SO Biological Psychiatry (2005), 57(8), 943-946 CODEN:

BIPOBF; ISSN: 0006-3223

PB Elsevier Inc.

DT Journal

LA English

AB Background: Recently a novel ***tryptophan***
 hydroxylase ***isoform*** (***TH2***) was
 identified and shown to be highly expressed in the central
 nervous system (CNS). Hormonal effects on ***TH2***
 mRNA expression in the rodent dorsal raphe nucleus (DRN) are
 unknown. Methods: In situ hybridization histochem. and real-
 time reverse transcriptase-polymerase chain reaction (RT-PCR)
 were used to assess the effects of dexamethasone or estradiol on
 TH2 mRNA levels in the DRN of C57/B6 mice. Results:
 Dexamethasone reduced ***TH2*** mRNA levels in the DRN
 of both ovx female and intact male mice. Redn. of ***TH2***
 mRNA in the DRN was blocked by co-administration of
 mifepristone. Estradiol had no detectable effect on
 TH2 mRNA levels in the DRN. Conclusions:
 TH2 mRNA is regulated by glucocorticoids but not
 estradiol in the mouse DRN. Glucocorticoid-mediated redn. of
 TH2 message may have relevance to the etiol. of major
 depression, psychotic major depression in particular, where
 elevated glucocorticoids are one hallmark of the disease.

L3 ANSWER 5 OF 28 CAPLUS COUNTRY GHT 2005 ACS ON STN
 AN 2005.307631 CAPLUS

TI Monoamine oxidase A and tryptophan hydroxylase gene
 polymorphisms: are they associated with bipolar disorder?

AU Preisig, Martin; Ferrero, Francois; Malafosse, Alain

CS University Department of Adult Psychiatry, Lausanne, Switz.
SO American Journal of Pharmacogenomics (2005), 5(1), 45-52
CODEN: AJPMG; ISSN: 1175-2203
PB Adis International Ltd.
DT Journal

LA English
AB Most of the candidate gene studies in bipolar disorder have focused on the major neurotransmitter systems that are influenced by drugs used in the treatment of this disorder. The monoamine oxidase A (MAOA) and the tryptophan hydroxylase (TPH1, ***TPH2***) genes are two of the candidates that have been tested in a series of association studies using unrelated or family-based controls. This review summarizes the existing association studies regarding these genes. Most of these studies were based on the unrelated case-control design with samples of 50 to 600 subjects. Regarding MAOA, three meta-analyses with partially overlapping samples supported a modest effect of this gene in bipolar disorder in female Caucasians. However, as several studies could not replicate these findings, more work is necessary to demonstrate unequivocally the involvement of MAOA in bipolar disorder and establish the biological mechanism underlying the genetic association. With respect to TPH1 and ***TPH2***, the majority of studies did not provide evidence for an association between these genes and bipolar disorder. The genes are more likely to be related to suicidal behavior than to bipolar disorder.

RE QNT 63 THERE ARE 63 Q TIED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 6 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN AN 2005:91263 CAPLUS
DN 142:311855

TI Different properties of the central and peripheral forms of human tryptophan hydroxylase
AU McKinney, Jeffrey, Knappskog, Per M.; Haavik, Jan
CS Department of Biomedicine, Section of Biochemistry and Molecular Biology, University of Bergen, Bergen, Norway
SO Journal of Neurochemistry (2005), 92(2), 311-320 CODEN: JONRA9; ISSN: 0022-3042

PB Blackwell Publishing Ltd.
DT Journal
LA English
AB Tryptophan hydroxylase (TPH) catalyzes the rate-limiting reaction in the biosynthesis of serotonin. In humans, two different TPH genes exist, located on chromosomes 11 and 12, respectively, encoding two enzymes (TPH1 and ***TPH2***) with an overall sequence identity of 71%. The authors have expressed both enzymes as various fusion proteins in *Escherichia coli* and using an in vitro transcription/translation system, and compared their solubility and kinetic properties. ***TPH2*** is more soluble than TPH1, has a higher molecular weight, and different kinetic properties, including a lower catalytic efficiency towards phenylalanine than TPH1. Both enzymes are phosphorylated by cAMP-dependent protein kinase A. ***TPH2*** was phosphorylated at Ser-19, a phosphorylation site not present in TPH1. The differences between TPH1 and ***TPH2*** have important implications for the regulation of serotonin production in the brain and the periphery and may provide an explanation for some of the diverging results reported for TPH from different sources in the past.

RE QNT 43 THERE ARE 43 Q TIED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 7 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN AN 2004:885157 CAPLUS

DN 142:111677
TI Support for the involvement of ***TPH2*** gene in affective disorders
AU Harvey, M.; Shink, E.; Tremblay, M.; Gagne, B.; Raymond, C.; Labbe, M.; Walther, D. J.; Bader, M.; Barden, N.
CS Neuroscience, CHUL Research Center, Ste-Foy, QC, Can.
SO Molecular Psychiatry (2004), 9(11), 980-981 CODEN: MOPSPQ; ISSN: 1359-4184
PB Nature Publishing Group
DT Journal
LA English
AB The tryptophan hydroxylase 2 (***TPH2***) gene was examined as a putative candidate gene using a SNP-based association study involving 213 individuals with bipolar disorder and 214 control subjects. Five out of the eight studied haplotypes represent 90 and 93% of haplotypes found in case and control groups, respectively. The data obtained provide support to previous results that suggest the existence of an affective disorder-associated haplotype in the ***TPH2*** gene.
RE QNT 9 THERE ARE 9 Q TIED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 8 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN AN 2004:885151 CAPLUS
DN 142:36363

TI SNP and haplotype analysis of a novel ***tryptophan*** hydroxylase*** isoform*** (***TPH2***) gene provide evidence for association with major depression
AU Zili, P.; Baghai, T. C.; Zwaninger, T.; Schuele, C.; Eser, D.; Rupprecht, R.; Moeller, H.-J.; Bondy, B.; Ackenheil, M.
CS Psychiatric Hospital, Ludwig-Maximilians-University, Munich, Munich, D-80336, Germany
SO Molecular Psychiatry (2004), 9(11), 1030-1036 CODEN: MOPSPQ; ISSN: 1359-4184
PB Nature Publishing Group
DT Journal
LA English

AB Tryptophan hydroxylase (TPH), being the rate-limiting enzyme in the biosynthesis of serotonin, plays a major role as a candidate gene in several psychiatric disorders. Recently, a second TPH isoform (***TPH2***) was identified in mice, which was exclusively present in the brain. In a previous post-mortem study of our own group, we could demonstrate that ***TPH2*** is also expressed in the human brain, but not in peripheral tissues. This is the first report of an association study between polymorphisms in the ***TPH2*** gene and major depression (MD). We performed single-nucleotide polymorphism (SNP), haplotype and linkage disequilibrium studies on 300 depressed patients and 265 healthy controls with 10 SNPs in the ***TPH2*** gene. Significant association was detected between one SNP (P=0.0012, global P=0.0051) and MD. Haplotype analysis produced additional support for association (P=0.0001, global P=0.0001). Our findings provide evidence for an involvement of genetic variants of the ***TPH2*** gene in the pathogenesis of MD and might be a hint on the repeatedly discussed duality of the serotonergic system. These results may open up new research strategies for the analysis of the observed disturbances in the serotonergic system in patients suffering from several other psychiatric disorders.

RE QNT 30 THERE ARE 30 Q TIED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 9 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN AN 2004:832118 CAPLUS
DN 142:4738

TI Single nucleotide polymorphism and haplotype analysis of a novel ***tryptophan*** **hydroxylase***

isoform (***TPH2***) gene in suicide victims
AU Zill, Peter; Buettner, Andreas; Eisenmenger, Wolfgang;
Moeller, Hans-Juergen; Bondy, Brigitta; Ackenheil, Manfred
CS Psychiatric Hospital, Ludwig-Maximilians-University, Munich,
Germany

SO Biological Psychiatry (2004), 56(8), 581-586 CODEN:
BIOPSF; ISSN: 0006-3223

PB Elsevier Inc.

DT Journal

LA English

AB Serotonin hydroxylase, the rate-limiting enzyme in the biosynthesis of serotonin, represents a major candidate in numerous genetic association analyses of suicidal behavior; however, the results are so far inconclusive. Recently, a second Trp hydroxylase isoform (***TPH2***) was identified in mice, which was exclusively present in the brain. In a previous postmortem study of our own group, the authors could demonstrate that ***TPH2*** is also expressed in the human brain but not in peripheral tissues. We performed single nucleotide polymorphisms, haplotypes, and linkage disequilibrium studies on 263 suicide victims and 266 healthy control subjects with 10 single nucleotide polymorphisms in the ***TPH2*** gene. Significant association ($p = .004$, global $p = .01$) and suicide. Additionally, haplotype analysis also produced support for association ($p < .0001$, global $p = .0001$). This is the first report about an association between ***TPH2*** gene polymorphisms and completed suicide. These findings provide evidence for an involvement of genetic variants in the ***TPH2*** gene in suicidal behavior. These results might open up new research strategies for the analysis of the observed disturbances in the serotonergic system in several other psychiatric disorders.

RE QNT 38 THERE ARE 38 Q/TED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q/TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 10 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN
AN 2004:786525 CAPLUS
DN 142:91296

TI Analysis of the novel ***TPH2*** gene in bipolar disorder and suicidality

AU De Luca, V.; Mueller, D. J.; Tharmalingam, S.; King, N.;
Kennedy, J. L.

CS Neurogenetics Section, Clarke Site, Centre for Addiction and
Mental Health, Department of Psychiatry, University of Toronto,
Toronto, ON, Can.

SO Molecular Psychiatry (2004), 9(10), 896-897 CODEN:
MOPSPF; ISSN: 1359-4184

PB Nature Publishing Group

DT Journal

LA English

AB The putative role of the tryptophan hydroxylase 2 (***TPH2***) gene in suicide attempters was studied using a large and well-characterized sample of patients with bipolar disorder. The study population consisted of 336 bipolar patients from 305 families who were recruited in the Toronto area. Among the bipolar patients, 267 patients had history of suicide ideas or attempt. The genotypes of ***TPH2*** hCV245410, hCV8376173, and rs1487280 polymorphisms were determined by Taqman assay. Allelic and haplotype transmission tests in suicide attempters were performed using TDT and TRANSMIT. When the suicide behavior was analyzed as a quantitative trait by the family-based association test, no significant differences were found for the three polymorphisms. All the eight possible haplotypes showed no significant results when weighted on the suicide behavior as

the quantitative trait. In bipolar patients, the TDT did not show significant bias for hCV245410, hCV8376173, and rs1487280.
RE QNT 13 THERE ARE 13 Q/TED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q/TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 11 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN
AN 2004:699213 CAPLUS

DN 142:107190

TI Investigation of serotonin-related genes in antidepressant response

AU Peters, E. J.; Sager, S. L.; McGrath, P. J.; Knowles, J. A.;
Hamilton, S. P.

CS Department of Psychiatry, University of California, San
Francisco, CA, USA

SO Molecular Psychiatry (2004), 9(9), 879-889 CODEN:
MOPSPF; ISSN: 1359-4184

PB Nature Publishing Group

DT Journal

LA English

AB In this study, we sought out to test the hypothesis that genetic factors may influence antidepressant response to fluoxetine. The investigation focused on seven candidate genes in the serotonergic pathway involved in the synthesis, transport, recognition, and degradation of serotonin. Our clinical sample consisted of 96 subjects with unipolar major depression treated with fluoxetine with response variables assessed after a 12-week trial. Patient data were also collected to investigate the pattern of drug response. Using a high-throughput single-nucleotide polymorphism (SNP) genotyping platform and capillary electrophoresis, we genotyped patients at 110 SNPs and four repeat polymorphisms located in seven candidate genes (HTR1A, HTR2A, HTR2C, MAOA, SLC6A4, TPH1, and ***TPH2***). Statistical tests performed included single-locus and haplotype association tests, and linkage disequilibrium (LD) estimation. Little evidence of population stratification was observed in the sample with 20 random SNPs using a genomic control procedure. Our most intriguing result involved three SNPs in the TPH1 gene and one SNP in the SLC6A4 gene, which showed significant single-locus association when response to fluoxetine was compared to nonresponse ($P = 0.02$ - 0.04). All odds ratios indicated an increased risk of not responding to fluoxetine. In the specific response vs. nonspecific and nonresponse comparison, three SNPs in the ***TPH2*** gene ($P = 0.02$ - 0.04) were positively associated, and one SNP in the HTR2A gene ($P = 0.02$) was negatively associated. When comparing specific response to nonspecific response, we found significant negative associations in three SNPs in the HTR2A gene ($P = 0.001$ - 0.03) and two SNPs in the MAOA gene ($P = 0.03$ - 0.05). We observed, although strong LD, in each gene and unexpectedly low number of estimated haplotypes, formed from tagged SNPs. Significant haplotype associations were found in all but the HTR1A and HTR2C genes. Although these data should be interpreted cautiously due to the small sample size, these results implicate TPH1 and SLC6A4 in general response, and HTR2A, ***TPH2***, and MAOA in the specificity of response to fluoxetine. Intriguingly, we observe that a number of the less frequent alleles of many of the SNP markers were associated with the nonresponse and nonspecific phenotypes.

RE QNT 61 THERE ARE 61 Q/TED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q/TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 12 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN
AN 2004:568127 CAPLUS

DN 141:185419

TI Brevia: Tryptophan hydroxylase-2 controls brain serotonin synthesis

AU Zhang, Xiaodong; Beaulieu, Jean-Martin; Sotnikova, Tatyana D.; Gainetdinov, Raul R.; Caron, Marc G.
CS Howard Hughes Med. Inst. Lab., Dep. Cell Biology and Center Models of Human Disease, Inst. Genome Sciences and Policy, Univ. Med. Center, Durham, NC, 27710, USA
SO Science (Washington, DC, United States) (2004), 305(5681), 217 CODEN: SSCI; ISSN: 0036-8075
PB American Association for the Advancement of Science
DT Journal
LA English
AB A minireview discussing the function of tryptophan hydroxylase-2 on controlling synthesis of brain serotonin.
RE QNT 10 THERE ARE 10 Q-TED REFERENCES AVAILABLE FOR THIS RECORD ALL Q-TATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 13 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 2004:487175 CAPLUS DN 141:289728

TI Transmission disequilibrium studies in children and adolescents with obsessive-compulsive disorders pertaining to polymorphisms of genes of the serotonergic pathway
AU Walitza, S.; Wewetzer, C.; Gerlach, M.; Klampt, K.; Geller, F.; Barth, N.; Hahn, F.; Herpertz-Dahlmann, B.; Goessler, M.; Fleischaker, G.; Schulz, E.; Hebebrand, J.; Warnke, A.; Hinney, A.

CS Department of Child and Adolescent Psychiatry, Julius-Maximilians-University, Würzburg, Germany
SO Journal of Neural Transmission (2004), 111(7), 817-825 CODEN: JNTRF3; ISSN: 0300-9564
PB Springer-Verlag Wien
DT Journal
LA English

AB Pharmacol. and challenge study data showed an involvement of the serotonergic system in the development of obsessive-compulsive disorder (OCD). We studied transmission disequilibrium of polymorphisms in three candidate genes of the serotonergic pathway in 64 trios comprising patients with early onset OCD and both of their parents. Polymorphisms of the following genes were studied: tryptophan hydroxylase 1 (rs1800532), serotonin transporter (polymorphism in the promoter region: 5-HTTLPR) and the serotonin 1B receptor (rs6296). This is, to our knowledge, one of the first family based assoc. studies pertaining to children and adolescents with OCD. We did not detect transmission disequilibrium of the investigated polymorphisms in OCD. Hence, these polymorphisms do not play a major role in the genetic predisposition to early onset OCD.
RE QNT 38 THERE ARE 38 Q-TED REFERENCES AVAILABLE FOR THIS RECORD ALL Q-TATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 14 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 2004:479424 CAPLUS DN 141:120812

TI Diurnal rhythms of tryptophan hydroxylase 1 and 2 mRNA expression in the rat retina
AU Liang, Jian; Wessel, James H., III; Iuvone, P. Michael; Tosini, Gianluca; Fukuhara, Osaki
CS Neuroscience Institute and NSF Center for Behavioral Neuroscience, Morehouse School of Medicine, Atlanta, GA, 30310-1495, USA
SO NeuroReport (2004), 15(9), 1497-1500 CODEN: NERPEZ; ISSN: 0959-4965
PB Lippincott Williams & Wilkins
DT Journal
LA English

AB Tryptophan hydroxylase is the first of four enzymes in the melatonin biosynthetic pathway. Recent studies have shown that there are two genes, Tph1 and ***Tph2***, that encode tryptophan hydroxylase in mammals. In this study, we investigated which of the two genes is expressed in the rat retina. To that end, we measured Tph1 (classical Tph) and ***Tph2*** mRNA levels using real-time quant. RT-PCR in the retina. Our data demonstrate that Tph1 mRNA is the prevalent form expressed in the retina; ***Tph2*** mRNA is also present but the level is very low. We also measured Tph1 expression levels in the outer nuclear layer, inner nuclear layer, and ganglion cell layer by combining laser capture microdissection and real-time RT-PCR. Tph1 mRNA is more abundant in the photoreceptors of the outer nuclear layer than in the inner nuclear layer or ganglion cell layer. Tph1 and ***Tph2*** transcripts showed robust diurnal rhythms of abundance, with highest levels at night. Our results support the hypothesis that Tph1 is involved in melatonin synthesis in retinal photoreceptor cells.
RE QNT 14 THERE ARE 14 Q-TED REFERENCES AVAILABLE FOR THIS RECORD ALL Q-TATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 15 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 2004:460775 CAPLUS DN 141:104429

TI Abnormal cardiac activity in mice in the absence of peripheral serotonin synthesis
AU Cote, Francine; Rigny, Cecile; Mallet, Jacques; Vojdani, Guilain
CS Laboratoire de Genetique Moleculaire de la Neurotransmission et des Processus Neurodegeneratifs, CNRS UMR7091, Hopital de la Pitie-Salpetriere, Paris, F-75013, Fr.
SO Journal de la Société de Biologie (2004), 198(1), 7-17 CODEN: JSBPGF; ISSN: 1295-0661
PB Masson Editeur
DT Journal
LA French

AB Serotonin (5-HT) controls a wide range of biol. functions. In the brain, its implication as a neurotransmitter and in the control of behavioral traits has been largely documented. At the periphery, its modulatory role in physiol. processes, such as the cardiovascular function, is still poorly understood. The rate limiting enzyme of 5-HT synthesis, tryptophan hydroxylase (TPH), is encoded by two genes: the well characterized TPH1 gene and a recently identified ***TPH2*** gene. Based on the study of a mutant mouse in which the TPH1 gene has been inactivated by replacement of the beta-galactosidase gene, we established that the neuronal ***TPH2*** is expressed in neurons of the raphe nuclei and of the myenteric plexus, whereas the non-neuronal TPH1, as detected by beta-galactosidase expression, is expressed in the pineal gland and the enterochromaffin cells. Anal. examn. of the mutant mice revealed larger heart sizes as compared to wild-type. Histol. investigations indicated that the primary structure of the heart muscle is not affected. Hemodynamic analyses in mutant animals demonstrated abnormal cardiac activity which ultimately leads to heart failure. This is the first report linking loss of TPH1 gene expression, and thus of peripheral 5-HT, to a cardiac dysfunction phenotype. The TPH1-/- mutant may be a valuable model for investigating cardiovascular dysfunction such as those obsd. in human heart failure.
RE QNT 44 THERE ARE 44 Q-TED REFERENCES AVAILABLE FOR THIS RECORD ALL Q-TATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 16 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN

AN 2004:419767 CAPLUS
DN 141:121683

TI Regional mRNA expression of a second ***tryptophan***
hydroxylase ***isoform*** in postmortem tissue

samples of two human brains

AU Zili, Peter; Buttner, Andreas; Eisenmenger, Wolfgang;

Bondy, Brigitta; Ackenheil, Manfred

CS Department of Psychiatry, Ludwig-Maximilians-University
Munich, Munich, D-80336, Germany

SO European Neuropsychopharmacology (2004), 14(4), 282-
284 CODEN: EUPNEB; ISSN: 0924-977X

PB Elsevier Science B.V.

DT Journal

LA English

AB Tryptophan hydroxylase (TH) as rate limiting enzyme in the
biosynthesis of serotonin plays a major role as candidate gene in
several psychiatric disorders. Recently a second TPH isoform (
TH2) was identified in mice, which was exclusively
expressed in the brain. We investigated whether the mRNA of
the human homolog of this new ***TH2*** isoform is

expressed in the human brain but not in peripheral tissues. The
study was performed with postmortem specimen obtained from
two subjects who died on cardiovascular failure. ***TH2***
mRNA levels were detd. by quant. real time RT-PCR

TH2 mRNA was exclusively present in the human
brains but not in the investigated peripheral tissues. Our finding
may open up new research strategies for the anal. of the
repeatedly obsd. disturbances in the serotonergic system in
patients suffering from several psychiatric disorders.

RE QNT 14 THERE ARE 14 Q TIED REFERENCES AVAILABE
FOR THIS RECORD ALL Q TATIONS AVAILABE IN THE RE
FORMAT

L3 ANSWER 17 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN
AN 2004:160595 CAPLUS
DN 140:315537

TI Serotonin regulates mammary gland development via an
autocrine-paracrine loop

AU Matsuda, Manabu; Imaoka, Tatsuhiro; Vomacka, Archie J.;
Gudelsky, Gary A.; Hou, Zhaoxuan; Mistry, Meenakshi; Bailey,
Jason P.; Newport, Kathryn M.; Walther, Diego J.; Bader, Michael;
Horseman, Nelson D.

CS Department of Molecular and Cellular Physiology, University
of Cincinnati, Cincinnati, OH, 45221, USA

SO Developmental Cell (2004), 6(2), 193-203 CODEN: DCEEBE;
ISSN: 1534-5807

PB Cell Press

DT Journal

LA English

AB Mammary gland development is controlled by a dynamic
interplay between endocrine hormones and locally produced
factors. Biogenic monoamines (serotonin, dopamine,
norepinephrine, and others) are an important class of
bioregulatory mol. that have not been shown to participate in
mammary development. Mammary glands stimulated by
prolactin (PRL) express genes essential for serotonin biosynthesis
(tryptophan hydroxylase [TH] and arom. amine decarboxylase).
TH mRNA was elevated during pregnancy and lactation, and
serotonin was detected in the mammary epithelium and in milk.
THP was induced by PRL in mammosphere cultures and by milk
stasis in nursing dams, suggesting that the gene is controlled by
milk filling in the alveoli. Serotonin suppressed beta-casein
gene expression and caused shrinkage of mammary alveoli.
Conversely, THP1 gene disruption or antiserotonergic drugs
resulted in enhanced secretory features and alveolar dilation.
Thus, autocrine-paracrine serotonin signaling is an important
regulator of mammary homeostasis and early involution.

RE QNT 34 THERE ARE 34 Q TIED REFERENCES AVAILABE
FOR THIS RECORD ALL Q TATIONS AVAILABE IN THE RE
FORMAT

L3 ANSWER 18 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN
AN 2004:111240 CAPLUS
DN 140:336184

TI Robust and tissue-specific expression of ***TH2***
versus THP1 in rat raphe and pineal gland

AU Patel, Pares D.; Pontrello, Crystal; Burke, Sharon
CS University of Michigan Medical Center, Ann Arbor, MI, USA

SO Biological Psychiatry (2004), 55(4), 428-433 CODEN:
BIOCBF; ISSN: 0006-3223

PB Elsevier Inc.

DT Journal

LA English

AB Background: Regulation of raphe serotonergic cells is
fundamental to the prevailing hypothesis of major depressive
pathophysiol. Tryptophan hydroxylase (THP) is the rate-limiting
enzyme in serotonin biosynthesis, but brainstem THP mRNA
expression has been difficult to measure and study. Recently, a
novel paralog of THP, ***TH2*** (or neuronal THP), was
described, but its anat. expression is unknown. Methods: An in
situ hybridization histochem. survey was conducted across
Sprague-Dawley rat brain for THP1 and ***TH2*** mRNA.
Semiquant. techniques were used to est. relative mRNA levels in
individual cells. Results: Almost exclusively, ***TH2***
mRNA is expressed in raphe, in a pattern overlapping the histol.
defined raphe nuclei. In sharp contrast, THP1 (the previously
known THP) is expressed predominantly in pineal gland. There is
no appreciable overlap in the expression of these paralogs. The
level of ***TH2*** mRNA expression in individual raphe cells
is approx. 2.5-fold greater than the level of THP1 expression in
pinealocytes. Conclusions: ***TH2*** mRNA has an anat.
expression pattern consistent with brainstem raphe nuclei and is
likely to be the gene giving rise to the majority of THP activity in
these cells. The robust expression of ***TH2*** in
regulation of raphe serotonin biosynthesis.

RE QNT 27 THERE ARE 27 Q TIED REFERENCES AVAILABE
FOR THIS RECORD ALL Q TATIONS AVAILABE IN THE RE
FORMAT

L3 ANSWER 19 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN
AN 2003:917153 CAPLUS
DN 140:53832

TI Disruption of the nonneuronal tph1 gene demonstrates the
importance of peripheral serotonin in cardiac function

AU Cote, Francine; Thevenot, Bienne; Fligny, Cecile; Fromes,
Yves; Darnon, Michele; Ripoche, Marie-anne; Bayard, Blisa;
Hannou, Naima; Saurin, Françoise; Lechat, Philippe; Dandolo,
Luiza; Hamon, Michel; Mallet, Jacques; Vojdani, Guilain
CS Laboratoire de Genetique Molculaire de la
Neurotransmission, Centre National de la Recherche Scientifique,
Unité Mixte de Recherche 7091 et Institut Fédératif de Recherche
70 (Neuroscience), Bâtiment CERVI, Hôpital de la Pitié-
Salpêtrière, Paris, 75013, Fr.

SO Proceedings of the National Academy of Sciences of the
United States of America (2003), 100(23), 13525-13530 CODEN:
PNAS66; ISSN: 0027-8424

PB National Academy of Sciences

DT Journal

LA English

AB Serotonin (5-HT) controls a wide range of biol. functions. In
the brain, its implication as a neurotransmitter and in the control
of behavioral traits has been largely documented. At the
periphery, its modulatory role in physiol. processes, such as the

cardiovascular function, is still poorly understood. The rate-limiting enzyme of 5-HT synthesis, tryptophan hydroxylase (TPH), is encoded by two genes, the well characterized tph1 gene and a recently identified "tph2" gene. In this article, based on the study of a mutant mouse in which the tph1 gene has been inactivated by replacement with the .beta.-galactosidase gene, the authors establish that the neuronal "tph2" gene is expressed in neurons of the raphe nuclei and of the myenteric plexus, whereas the nonneuronal tph1, as detected by .beta.-galactosidase expression, is in the pineal gland and the enterochromaffin cells. Anat. examn. of the mutant mice revealed larger heart sizes than in wild-type mice. Histol. investigation indicates that the primary structure of the heart muscle is not affected. Hemodynamic analyses demonstrate abnormal cardiac activity, which ultimately leads to heart failure of the mutant animals. This report links loss of tph1 gene expression, and thus of peripheral 5-HT, to a cardiac dysfunction phenotype. The tph1-/- mutant may be valuable for investigating cardiovascular dysfunction obsd. in heart failure in humans.
RE QNT 53 THERE ARE 53 Q TIED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 20 OF 28 CAPLUS COPYRHT 2005 ACS ON STN
AN 2003:808741 CAPLUS
DN 140:53582
TI A unique central "tryptophan" "hydroxylase"
"isoform"

AU Walther, Diego J.; Bader, Michael
CS Max Delbrück Center for Molecular Medicine (MDG), Berlin, D-13092, Germany
SO Biochemical Pharmacology (2003), 66(9), 1673-1680
CODEN: BCPAC6; ISSN: 0006-2952
PB Elsevier Science B.V.,
DT Journal; General Review
LA English
AB A review. Serotonin (5-hydroxytryptophan, 5-HT) is a neurotransmitter synthesized in the raphe nuclei of the brain stem and involved in the central control of food intake, sleep, and mood. Accordingly, dysfunction of the serotonin system has been implicated in the pathogenesis of psychiatric diseases. At the same time, serotonin is a peripheral hormone produced mainly by enterochromaffin cells in the intestine and stored in platelets, where it is involved in vasoconstriction, hemostasis, and the control of immune responses. Moreover, serotonin is a precursor for melatonin and is therefore synthesized in high amounts in the pineal gland. Tryptophan hydroxylase (TPH) catalyzes the rate limiting step in 5-HT synthesis. Until recently, only one gene encoding TPH was described for vertebrates. By gene targeting, we functionally ablated this gene in mice. To our surprise, the resulting animals, although being deficient for serotonin in the periphery and in the pineal gland, exhibited close to normal levels of 5-HT in the brain stem. This led us to the detection of a second TPH gene in the genome of humans, mice, and rats, called "TPH2". This gene is predominantly expressed in the brain stem, while the classical TPH gene, now called TPH1, is expressed in the gut, pineal gland, spleen, and thymus. These findings clarify puzzling data, which have been collected over the last decades about partially purified TPH proteins with different characteristics and justify a new concept of the serotonin system. In fact, there are two serotonin systems in vertebrates, independently regulated and with distinct functions.

RE QNT 93 THERE ARE 93 Q TIED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 21 OF 28 CAPLUS COPYRHT 2005 ACS ON STN
AN 2003:692174 CAPLUS
DN 139:289470
TI Comparison of circadian expression of "tryptophan" "hydroxylase" "isoform" mRNAs in the rat pineal gland using real-time PCR
AU Sugden, David
CS Centre for Reproduction, Endocrinology and Diabetes, School of Biomedical Sciences, Kings College London, London, UK
SO Journal of Neurochemistry (2003), 86(5), 1308-1311
CODEN: JONRA9; ISSN: 0022-3042
PB Blackwell Publishing Ltd.
DT Journal
LA English

AB A second gene encoding a functional tryptophan hydroxylase activity has recently been described ("TPH2"), which is expressed abundantly in brainstem, the primary site of serotonergic neurons in the CNS. As serotonin (5-HT) has an important role as a precursor of the nocturnal synthesis of the pineal gland hormone, melatonin, it was of interest to det. the relative expression of TPH1 and 2 mRNA in the rat pineal during the light/dark (LD) cycle using sensitive real-time RT-PCR assays which were developed for each TPH isoform. TPH1 mRNA expression was 105-fold more abundant in rat pineal than "TPH2", and showed a significant, approx. 4-fold nocturnal increase in expression which may contribute to the previously described nocturnal increase in pineal tryptophan hydroxylase activity. "TPH2" expression within the gland showed no significant variation with time of day and was very low (approx. 300 copies/gland) indicating expression in the small proportion of "non-pinealocyte" cells in the gland.
RE QNT 28 THERE ARE 28 Q TIED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 22 OF 28 CAPLUS COPYRHT 2005 ACS ON STN
AN 2003:384284 CAPLUS
DN 138:40160
TI Elisabeth-Gatteff prize of GFG 2001: Serotonin biosynthesis in the central nervous system will be rate-definably catalyzed by a neuron-specific Trp-hydroxylase-isozyme
AU Walther, Diego J.
CS Dept. of Genetics, Bioinformatics and Structural Biology, Max Delbrück Center for Molecular Medicine, Berlin, Germany
SO BioSpectrum (2003), 9(2), 184-186 CODEN: BOSFPD; ISSN: 0947-0867
PB Spektrum Akademischer Verlag
DT Journal
LA German

AB The work of the winners of the Elisabeth-Gatteff-Preis 2001 Diego J. Walther et al. is presented which is concerned with the discovery of a "tryptophan" "hydroxylase" "isoform" in the brain catalyzing serotonin biosynthesis in the central nervous system.
RE QNT 21 THERE ARE 21 Q TIED REFERENCES AVAILABLE
FOR THIS RECORD ALL Q TATIONS AVAILABLE IN THE RE
FORMAT

L3 ANSWER 23 OF 28 CAPLUS COPYRHT 2005 ACS ON STN
AN 2003:341202 CAPLUS
DN 140:89687
TI Knockout mouse points to second form of tryptophan hydroxylase
AU Veenstra-VanderWeele, Jeremy; Cook, Edwin H., Jr.
CS Department of Psychiatry, The University of Chicago, Chicago, IL 60637, USA

SO Molecular Interventions (2003), 3(2), 72-75 CODEN: MIONAR; ISSN: 1534-0384

SB American Society for Pharmacology and Experimental Therapeutics

DT Journal; General Review

LA English

AB A review describes the identification of two tryptophan hydroxylase (TPH) mRNA species arising from different promoters and having different translational efficiency. The expression of more efficiently translated isoform is increased in response to stress. The study by Walther et al. (2003) demonstrated that a Tph1 knockout mouse failed to generate the expected phenotype, in which Tph1-deficient mice continued to produce 5-hydroxytryptamine in the brain, but had almost no detectable serotonin in the duodenum, and none in the whole blood. The study found that the antibodies commonly used to identify TPH cross-reacted to both Tph1 and ****Tph2*** gene products. RE QNT 10* THERE ARE 10 QTED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAIL ABLE IN THE RE FORMAT

L3 ANSWER 24 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 1999:903198 CAPLUS DN 138:331851

TI Synthesis of serotonin by a second ****tryptophan*** **hydroxylase*** **isoform***

AA Walther, Diego J.; Peter, Jens-Uwe; Bashammakh, Saleh; Hortnagl, Heide; Voits, Mechthild; Fink, Heidrun; Bader, Michael SC Max Delbrück Center Molecular Med. (MDC), Berlin, D-13092, Germany

SO Science (Washington, DC, United States) (2003), 299(5603), 76 CODEN: SCIEAS; ISSN: 0036-8075

SB American Association for the Advancement of Science

DT Journal

LA English

AB Serotonin (5-HT) is synthesized in two steps, with tryptophan hydroxylase (TPH) as the rate-limiting enzyme. To study the physiol. impact of the loss of 5-HT synthesis, the authors generated mice genetically deficient for TPH (Tph-1-). Tph-/- mice expressed normal amts. of 5-HT in classical serotonergic brain regions. However, Tph-/- mice lacked 5-HT in the periphery except for in the duodenum. Tph-/- mice exhibited no significant behavioral differences in elevated plus maze and hole board tests, which are indicative for 5-HT related behavior. Despite suggestions of a possible second TPH isoform mol. verification has been lacking. The authors therefore cloned and sequenced the TPH isoform (referred as ****Tph2***) (GenBank: AY090565) which was different from the known TPH (referred to as Tph1), Pah and Th of the mouse. Tph1 mRNA was detected in the duodenum, but not in the brain. In contrast ****Tph2*** was detected exclusively in the brain. In addn. the authors also cloned and sequenced the rat and human ****TPH2*** homologs (GenBank: AY098915 and AY098914). The discovered duality of the serotonin system in vertebrates may open up new avenues for specific therapeutic approaches exclusively affecting central or peripheral 5-HT actions.

L3 ANSWER 25 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 1999:754632 CAPLUS DN 132:152131

TI Silylcupration of (R)-2,2-Dimethyl-3-(tert-butoxycarbonyl)-4-ethynylloxazolidine: A Stereoselective Approach to the Synthesis of gamma-Silylated Saturated and Unsaturated alpha-Amino Acids

AU Reginato, Ganna; Mordini, Alessandro; Valacchi, Michela; Grandini, Bena

CS Dipartimento di Chimica Organica U. Schiff, Centro CNR Composti Biorcoidici, Florence, I-50121, Italy

SO Journal of Organic Chemistry (1999), 64(25), 9211-9216 CODEN: JOCEAH; ISSN: 0022-3263

SB American Chemical Society

DT Journal

LA English

CS CASREACT 132:152131

AB Enantioselective synthesis of gamma-silylated amino acids is reported, using a four-step procedure based on the silylcupration of ethynylloxazolidine (I). Silylcuprates are highlighted as useful reagents to be employed with enantiomerically enriched substrates. Vinylsilanes [1]; SF3 S SMe3, SiPhMe2, or SiBu- ****TPH2***] are easily prep. and highlighted as useful intermediates to yield the final compds. after reduct., opening of the oxazolidine ring, and oxidn. Moreover, beta-, gamma-unsatd. amino acids are obtained as very interesting vinylglycine derivs. The capability of silicon-contg. amino acids to be incorporated into dipeptides is also shown.

RE QNT 34* THERE ARE 34 QTED REFERENCES AVAILABLE FOR THIS RECORD ALL Q TATIONS AVAIL ABLE IN THE RE FORMAT

L3 ANSWER 26 OF 28 CAPLUS COPYRIGT 2005 ACS ON STN AN 1999:298741 CAPLUS DN 131:38746

TI Inter- and Intramolecular Hydrogen-Bonding Interaction of Hydroxo Groups and Steric Effect of Alkyl Substituents on Pyrazolyl Rings in TrR Ligands: Synthesis and Structural Characterization of Chloro-, Acetylacetonato-, and Hydroxo Complexes of VO2+ with TrPr12 and TrMe2 Ligands

AU Kosugi, Masahiro; Hikichi, Shiro; Akita, Munetaka; Moro-oka, Yoshihiko

CS Research Laboratory of Resources Utilization, Tokyo Institute of Technology, Yokohama, 226-8503, Japan

SO Inorganic Chemistry (1999), 38(11), 2567-2578 CODEN: INOCAJ; ISSN: 0020-1669

SB American Chemical Society

DT Journal

LA English

AB Novel vanadyl (VO2+) chloro and hydroxo complexes with the hindered TrPr12 (hydrotris(3,5-diisopropyl-1-pyrazolyl)borate) and TrMe2 (hydrotris(3,5-dimethylpyrazolyl-1-pyrazolyl)borate) ligands were prep. and structurally characterized successfully. Ligand displacement of VO2(MeO)2(H2O) by TrR afforded octahedral chloro complexes, TrR(VO)Cl(X) (1: R = Pr12, X = TrPr12H; 2: R = Pr12, X = py; 3: R = Me2, X = NCOMe). Hydrolysis of the obtained chloro complexes yielded the corresponding hydroxo complexes 4, 5, and 7, but their structures were very unique and different from that of the previously reported dinuclear VO2+ bis(mu-hydroxo) complex with the less hindered ****TPH2*** (hydrotris(1-pyrazolyl)borate) ligand. For the TrPr12 complexes, the octahedral hydroxo-aqua complex, TrPr12V(O)(OH)(OH2) (4), and the trinuclear bis(mu-hydroxo)bis(mu-pyrazolate) complex, TrPr12V(O)(mu-OH)(mu-PrPr12)V(O)(mu-OH)(mu-PrPr12)V(O)TrPr12 (5), were isolated. The hydroxo-aqua complex 4 was dimerized through the internal hydrogen-bonding interaction between the hydroxo and aqua ligands forming the H3O2-bridging ligand. The trinuclear complex 5 consisted of two octahedral TrPr12V(O) fragments and a distorted trigonal-bipyramidal non-TrPr12V(O)-supported VO2+ center, sitting on the pseudo C2 symmetry axis, and was formed via coupling of 4 and the VO2+ pyrazolate species, resulting from partial decomp. of the chloro complexes during the hydrolysis. Steric repulsion of the bulky Pr groups in TrPr12 might hinder the formation of a dinuclear bis(mu-

hydroxo) complex like the ***Tph2*** and Tpm2 derivs. The dinuclear bis-(mu - hydroxo) complex with the Tpm2 ligand, (kappa 3-Tpm2)W(O)(mu - OH)2W(O)(kappa 2-Tpm2) (7), consisted of syn-arranged VO fragments, having the different coordination geometries of the vanadium centers (octahedron with kappa 3-Tpm2 and trigonal bipyramid with kappa 2-Tpm2). Intramol. hydrogen-bonding interaction between one of the two hydroxo groups and the noncoordinated pyrazolyl nitrogen atom in kappa 2-Tpm2 was obsd.
RE QUT 81 THERE ARE 81 QUTED REFERENCES AVAILABLE FOR THIS RECORD ALL QUTATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 27 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN
AN 1996:308899 CAPLUS
DN 124:336147
TI Genetic variability of tocopherol composition in sunflower seeds: as a basis of breeding for improved oil quality
AU Demurin, Y.; Gkoric, D.; Karlovic, D.
CS Rustovits Institute Oil Crops (VNIIMK), Krasnodar, 38, Russia
SO Plant Breeding (1996), 115(1), 33-36 CODEN: PLABED; ISSN: 0179-9541
PB Blackwell
DT Journal
LA English
AB The variability of seed tocopherol content in wild sunflower species, the expressivity of tph1 and ***tph2*** mutations in different lines and the oxidative stability of sunflower oil with altered tocopherol and fatty acid comp. were objectives of this research. Near-isogenic lines for three genes, i.e. Tph1, ***Tph2***, and QI, were developed and investigated. Tocopherol content was detd. with TLD and HPLC, as well as fatty acid comp. with GC of Me esters. Rancimat tests were used to est. the oxidative stability of the oil. The seed tocopherol compn. of wild sunflower species was shown to be uniform with a prevailing content of the alpha-homolog (90-99%). The genetic background of different near-isogenic lines was found to influence expressivity of mutations for tocopherol compn. High content of strong antioxidants, such as beta-, gamma-, and delta-tocopherols increased oil oxidative stability of linoleic and oleic types of oil by 1.2-3.0 times. The breeding model of sunflower hybrids should include antioxidant and vitamin parameters balanced for oils of different applications.

L3 ANSWER 28 OF 28 CAPLUS COPYRIGHT 2005 ACS ON STN
AN 1986:572614 CAPLUS
DN 105:172614
TI Tin-119 Moessbauer spectroscopic studies of the products of the reaction of triorganotin(IV) derivatives with 6-thiopurine
AU Barbieri, R.; Di Bianca, F.; Rivaola, E.; Huber, F.
CS Ist. Chim. Gen., Univ. Palermo, Palermo, I-90123, Italy
SO Inorganica Chimica Acta (1985), 108(3), 141-5 CODEN: ICHAAA; ISSN: 0020-1693
DT Journal
LA English
CS CASREACT 105:172614
AB A structural study of the products of the reaction of R3Sn(IV) derivs. (R = Me, Bu, Ph) with 6-thiopurine, 6-***TPH2***, and its sodium salt, 6-TPHNa, has been undertaken using Moessbauer spectroscopy and the point-charge model rationalization of the Moessbauer parameter nuclear quadrupole splitting. The synthetic reactions have been carried out at approx. 0, 20, and 50 degree. The Moessbauer spectra of the complexes R3Sn(6-TPH) are consistent with the occurrence of two distinct tin(IV) sites in samples prepd. at the lower temp., while one only site appears by increasing the temp. of the

reaction. Two tin sites constantly occur in the products of the reactions involving the R3Sn(IV) moiety; the stoichiometry is assumed to be (R3Sn)3(6-TPH)(6-TP) for the uniquely formed complex. Solid state polymeric structures with trigonal bipyramidal environments of the tin atoms and planar SnC3 skeletons have been proposed. The apical ligand atoms have been assumed to be N, S and N, N in the samples showing two individual tin(IV) sites, and N, N when a single site was present.

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L1 15 S
L2 (TRYPTOPHAN(W)HYDROXYLASE(W))SOFORM)/BI,AB
L3 23 S TPH2/BI,AB
L3 28 S L1 OR L2
=> log y
COST IN U.S. DOLLARS SINCE FILE
TOTAL ENTRY SESSION
FULL ESTIMATED COST 90.67 90.88

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE
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